

L3 ANSWER 1 OF 23 HCAPLUS COPYRIGHT 2000 ACS

AN 2000:246339 HCAPLUS

T1 Hydroxyurea and ***trimidox*** enhance the radiation effect in human pancreatic adenocarcinoma cells

AL1 Lyden, D.; Ahmed, N.; Hassan, H. T.

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SO Anticancer Res 1999; 19:133-138

CODEN: ANTIDH; ISSN: 0259-7535

PB International Institute of Anticancer Research

DT Journal

LA English

CC 8 (Radiation Biochemistry)

AB BACKGROUND: Pancreatic cancer remains the most lethal of all common human malignancies with a 5-yr survival rate of lower than 5%. Adjuvant and neoadjuvant preoperative and postoperative chemo-radiotherapy using 5-fluorouracil, have reduced local relapse rate and slightly increased the median survival. Testing new and more potent radiation-sensitizing drugs in human pancreatic cancer cells can provide the basis for a more effective chemo-radiation regimen and may consequently improve treatment outcome. MATERIALS & METHODS: The present study was performed to evaluate the efficacy of two potent ribonucleotide reductase (RR) inhibitors: hydroxyurea and ***trimidox*** in radio-sensitizing Panc-1 human pancreatic cancer cells in an attempt to identify a more effective chemo-radiotherapy regimen with minimal side effects. RESULTS: Treatment of Panc-1 cells with hydroxyurea or ***trimidox*** alone for 2 h was assoc. with a dose-dependent decrease in their cloning efficacy to similar extent. The IC50 of ***trimidox***, hydroxyurea or radiation alone were 2.5 + 0.3 .mu.M, 39.0 + 0.4 .mu.M and 3.2 + 0.2 Gy, resp. Treatment with 39.0 microM non-cytotoxic IC50 dose of hydroxyurea for two hours before or immediately after radiation reduced the IC50 of radiation to only 1.1 + 0.14 or 1.0 + 0.1 Gy, resp. Treatment with 2.5 microM non-cytotoxic IC50 dose of ***trimidox*** for two hours before or immediately after radiation reduced the IC50 of radiation to only 1.2 + 0.16 or 1.4 + 0.12 Gy, resp. The mean radiation enhancement ratios were 2.9 and 3.2 for hydroxyurea before and immediately after radiation. The greater radio-sensitizing effect of hydroxyurea compared to ***trimidox*** or gemcitabine could be due to its unique double action by synchronising the cancer cells into the radiosensitive G1/S border and inhibiting DNA damage repair. CONCLUSIONS: The present study demonstrates the superiority of hydroxyurea at non-cytotoxic doses compared to the other two recent RR inhibitors: gemcitabine and ***trimidox*** in radio-sensitizing human pancreatic cancer cells. Hydroxyurea combined with radiation has significantly improved progression-free survival of advanced cervical cancer and glioblastoma patients and showed clin. benefit in combination with other chemotherapy drugs in advanced pancreatic cancer. The present results suggest the clin. use of hydroxyurea as a radiosensitizer in both pre- and post-operative chemo-radiotherapy in pancreatic cancer patients. Given the demonstrated potent radio-sensitizing effect of hydroxyurea at non-cytotoxic doses when administered before or immediately after radiation and its low clin. toxicity, it could be feasible to administer hydroxyurea both before and after radiation in pancreatic cancer patients.

RE.CNT 35

RE

- (1) Andre, T; Presse Med 1999, V27, P539
- (2) Burris, H; Eur J Cancer 1997, V33(supplement 1), P518
- (3) Elford, H; Proceedings of AACR 1999, V40, P591
- (4) Gale, R; Blood 1998, V91, P1810 HCAPLUS
- (5) Gandhi, V; J Clin Oncol 1998, V16, P2321 HCAPLUS
- (6) Hall, E; Radiobiology for the Radiobiologist, Fourth edition 1994
- (7) Hassan, H; Anticancer Res 1991, V11, P481 MEDLINE
- (8) Hassan, H; Haematol Reviews Commun 1997, V7, P125
- (9) Hassan, H; J Cancer Res Clin Oncology 1991, V117, P227 MEDLINE
- (10) Hassan, H; J Interferon Cytokine Res 1995, V16, P139
- (11) Howara, T; Current Problems Cancer 1996, V20, P281
- (12) Ishikawa, O; Hepatogastroenterology 1998, V45, P644 MEDLINE
- (13) Iyama, E; Biochem Biophys Res Commun 1998, V247, P759 HCAPLUS
- (14) Kamthan, A; Clin Oncol 1997, V15, P2920 HCAPLUS
- (15) Kuo, M; Cancer Res 1998, V58, P2245 HCAPLUS
- (16) Kuo, M; The Cancer Journal from Scientific American 1997, V3, P163 MEDLINE
- (17) Lawrence, T; Int J Radiation Oncology Biology Physics 1996, V34, P867 HCAPLUS
- (18) Levin, V; Seminars Oncol 1992, V19(supplement 9), P34
- (19) Lofvenberg, E; Eur J Haematol 1988, V41, P375 MEDLINE
- (20) McGinn, C; Current Problems Cancer 1993, V17, P273 MEDLINE
- (21) McGinn, C; Journal of National Cancer Institute 1996, V88, P1193 HCAPLUS
- (22) Mornex, F; Cancer Radiotherapy 1998, V2, P696 MEDLINE
- (23) Pisters, P; J Clin Oncol 1998, V16, P3843 MEDLINE
- (24) Prott, F; Brit J Cancer 1997, V75, P597 HCAPLUS
- (25) Raymond, E; Eur J Cancer 1997, V33, P696 HCAPLUS
- (26) Rosier, J; Int J Radiation Biology 1999, V75, P245 HCAPLUS
- (27) Rubens, R; Brit J Cancer 1991, V64, P1187 MEDLINE
- (28) Schilsky, R; Seminars Oncol 1992, V19(supplement 9), P84
- (29) Shewach, D; Investigational New Drugs 1996, V1, P257

(30) Shtiz, P; J Clin Oncol 1997, V15, P924 MEDLINE

(31) Sporn, J; Drugs 1999, V57, P69 MEDLINE

(32) Stehman, P; J Clin Oncol 1993, V11, P1523 MEDLINE

(33) Stornicka, A; Cancer 1999, V85, P1261 HCAPLUS

(34) Szwarc, T; Cancer Chemotherapy and Pharmacology 1994, V34, P93 HCAPLUS

(35) Thomas, P; Hepatogastroenterology 1998, V45, P610 MEDLINE

L3 ANSWER 2 OF 23 HCAPLUS COPYRIGHT 2000 ACS

AN 2000:246339 HCAPLUS

T1 Enhancement of hemoglobin and F-cell production by targeting growth inhibition and differentiation of K562 cells with ribonucleotide reductase inhibitors (diox and ***trimidox***) in combination with streptozotocin

AL1 Iyama, W. E.; Adangish, S. E.; Finkeld, H.; Horiuchi, K.; Elford, H. L.; Ashkhar, T.; Turner, E. A.

CS Comprehensive Sickle Cell Center, Meharry Medical College, Nashville, TN, USA

SO Am. J. Hematol. (2000), 63(4), 176-183

CODEN: AJHEDD; ISSN: 0361-8609

PB Wiley-Liss, Inc.

DT Journal

LA English

CC 1 (Pharmacology)

AB Upon appropriate drug treatment, the human erythroleukemic K562 cells have been shown to produce Hb and F-cells. Fetal Hb (Hb F) inhibits the polymn. events of sickle Hb (Hb S), thereby ameliorating the clin. symptoms of sickle cell disease. Ribonucleotide reductase inhibitors (RRIs) have been shown to inhibit the growth of myeloid leukemia cells leading to the prodn. of Hb F upon differentiation. Of the RRIs currently in use, hydroxyurea is the most effective agent for Hb F induction. We have examd. the capacity of two novel RRIs, diox (DI) and ***trimidox*** (TRI), in combination with streptozotocin (STZ), to induce Hb and F-cell prodn. The K562 cells were cultured with different concns. of diox-STZ or ***trimidox*** -STZ at a fixed molar ratio of 3:1 and 1:5 for 96 h, resp. At pre-detd. time intervals, aliquots of cells were obtained and total Hb (benzidine pos.) levels, no. of F-cells, and Hb F were detd. by the differential staining technique, fetal Hb assay kit, and fluorescence cytometry resp. The effect of combined drug treatment on the growth of K562 cells was examd. by isobologram anal. Our results indicate that a synergistic growth-inhibitory differentiation effect occurred when diox or ***trimidox*** was used in combination with STZ on K562 cells. There was an increase in the no. of both benzidine-pos. normoblasts and F-cells, accompanied by morphol. appearances typical of erythroid maturation. On day 4, the no. of benzidine-pos. cells showed a 6-9-fold increase and the no. of F-cells was between 2.5- and 5.7-fold higher than the resp. controls. Based upon these results, treatment with a ribonucleotide reductase inhibitor, such as diox or ***trimidox***, in combination with STZ, might offer an addnl. promising option in sickle cell disease therapy.

RE.CNT 32

RE

- (1) Arwicz, A; J Surg Oncol 1979, V12, P267
- (2) Charache, S; Adv Pediatr 1990, V37, P1 MEDLINE
- (3) Charache, S; Blood 1992, V79, P2555 MEDLINE
- (4) Charache, S; N Engl J Med 1995, V332, P1317 MEDLINE
- (5) Chou, T; Adv Enzyme Regul 1984, V22, P27 HCAPLUS
- (6) DeSimone, J; Proc Natl Acad Sci USA 1982, V79, P4428 HCAPLUS
- (7) DeVita, V; Cancer, principles and practice of oncology 1989
- (8) Desesso, J; Teratology 1994, V49, P248 HCAPLUS
- (9) Ding, M; Ann Thorac Surg 1992, V53, P1091 MEDLINE
- (10) Elford, H; Biochem Biophys Res Commun 1968, V33, P129 HCAPLUS
- (11) Elford, H; Cancer Res 1979, V39, P844 HCAPLUS
- (12) Elford, H; Inhibitors of ribonucleoside diphosphate reductase activity 1989, P17
- (13) Fibach, E; Blood 1993, V81, P1630 HCAPLUS
- (14) Horiuchi, K; Biochem Biophys Res Commun 1995, V217, P924 HCAPLUS
- (15) Horiuchi, K; Cytometry 1995, V20, P261 MEDLINE
- (16) Horiuchi, K; Exp Hematol 1994, V22, P1058 MEDLINE
- (17) Iyama, E; J Chromatogr B 1998, V709, P119 HCAPLUS
- (18) Iyama, W; Biochem Biophys Res Commun 1998, V247, P759
- (19) Letvin, N; N Engl J Med 1984, V310, P869 HCAPLUS
- (20) McLeod, D; Blood 1974, V44, P517 MEDLINE
- (21) Moore, E; International encyclopedia of pharmacology and therapeutics 1989, P165
- (22) Noguchi, C; Blood 1981, V58, P1057 HCAPLUS
- (23) Pace, B; Am J Hematol 1994, V45, P136 HCAPLUS
- (24) Perrine, S; N Engl J Med 1993, V328, P81 HCAPLUS
- (25) Reilly, M; Exp Hematol 1994, V22, P501 HCAPLUS
- (26) Schechter, A; Molecular basis of blood diseases 1987, P187
- (27) Schein, P; Arch Intern Med 1973, V132, P555 MEDLINE
- (28) Szekeres, T; Cancer Chemother Pharmacol 1994, V34, P63 HCAPLUS
- (29) van't Ried, B; J Med Chem 1979, V22, P589
- (30) Veale, D; Br J Cancer 1988, V58, P70 HCAPLUS
- (31) Veith, R; N Engl J Med 1985, V313, P1571 MEDLINE
- (32) Zhao, K; Anticancer Res 1995, V15, P645 MEDLINE

L3 ANSWER 3 OF 23 HCAPLUS COPYRIGHT 2000 ACS

AN 2000:209828 HCAPLUS

DN 132:246339

T1 Antiviral drug compositions containing lithium salts

- (31) Mulvey, E; Proc Natl Acad Sci USA 1994, V91, P11087 HCAPLUS
 (32) Mayhew, C; Cell Mol Biol (Neuro-biogr) 1997, V43, P1019 HCAPLUS
 (33) Meyerhans, A; J Virol 1994, V68, P533 HCAPLUS
 (34) Mills, D; Antivir Res 1992, V17(suppl 1), P56
 (35) Mills, D; Proc Am Assoc Cancer Res 1992, V33, P799
 (36) Montaner, L; J Infect Dis 1997, V175, P601 HCAPLUS
 (37) Monte, B; Pharmacol Ther 1993, V27, P167 HCAPLUS
 (38) Nicot, E; Cell Tissue Res 1996, V268, P479 HCAPLUS
 (39) Neuman, E; Cancer Chemother Pharmacol 1997, V39, P234 HCAPLUS
 (40) Padilla, F; N Engl J Med 1993, V328, P833
 (41) Palmer, S; Antitumor Agents Chemother 1999, V43, P2046 HCAPLUS
 (42) Roca-Salvador, F; Pharmacotherapy 1999, V19, P196 HCAPLUS
 (43) Roberts, R; Br J Cancer 1991, V64, P1187 MEDLINE
 (44) Ruckenstein, O; AIDS 1994, V8, P771 HCAPLUS
 (45) Shaffer, R; Ann Intern Med 1998, V128, P906 HCAPLUS
 (46) Simonelli, C; J Acquir Immune Defic Syndr Hum Retrovirol 1996, V13, P462 MEDLINE
 (47) Smith, D; Cancer Chemother Pharmacol 1993, V33, P139 MEDLINE
 (48) Szekeres, T; Cancer Chemother Pharmacol 1994, V34, P63 HCAPLUS
 (49) Szekeres, T; Crit Rev Clin Lab Sci 1997, V34, P503 HCAPLUS
 (50) Ussery, M; Intersci Conf Antimicrob Agents Chemother 1996, P188
 (51) Uyei, E; J Pharmacol Exp Ther 1976, V198, P246 HCAPLUS
 (52) van T Riet, B; US 4942253 1990
 (53) van T Riet, B; J Med Chem 1979, V22, P589 HCAPLUS
 (54) Veale, D; Br J Cancer 1988, V58, P70 HCAPLUS
 (55) Veale, D; Cancer Chemother Pharmacol 1988, V21, P53 HCAPLUS
 (56) Wong, J; Proc Natl Acad Sci USA 1997, V94, P12574 HCAPLUS
 (57) Yarbro, J; Semin Oncol 1992, V19, P1 HCAPLUS

L3 ANSWER 5 OF 23 HCAPLUS COPYRIGHT 2000 ACS
 AN 1999-511044 HCAPLUS

DN 131:139487

TI Pharmaceutical compositions comprising PEG-asparaginase for the treatment of HIV infections

IN Avramis, Vassilios I.; Cohen, Lewis

PA Rhone-Poulenc Rorer Pharmaceuticals Inc., USA

SO PCT Int. Appl., 81 pp.

CODEN: PIXXD2

DT Patent

LA English

IC ICM A61K038-50

CC 1-5 (Pharmacology)

Section cross-reference(s): 63

FAN.CNT 1

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI WO 9939732	A1	19990812	WO 1999-US2480	19990209

W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE, GH, GM, HU, ID, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
 RW: GH, GM, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

AU 9926584 A1 19990823 AU 1999-26584 19990209

PRAI US 1998-PV74066 19980209

US 1998-74066 19980209

WO 1999-US2480 19990209

OS MARPAT 131:139487

AB A method of inhibiting or treating Human Immunodeficiency Virus (HIV) infection comprises administering to a patient in need thereof an effective amt. of a pharmaceutically acceptable compn. comprising a PEG-ASNase compd. and optionally at least one compd. selected from the group consisting of protease inhibitor compds., ribonucleotide reductase inhibitor compds. and HIV reverse transcriptase inhibitor compds.

ST PEG asparaginase HIV infection treatment

IT Antiviral agents

Drug delivery systems

Human immunodeficiency virus

Human immunodeficiency virus 1

(PEG-asparaginase, and combinations with other agents, for HIV infection treatment)

IT Amino acids, biological studies

RL: BOC (Biological occurrence); BPR (Biological process); BIOL (Biological study); OCCU (Occurrence); PROC (Process)

(PEG-asparaginase, and combinations with other agents, for HIV infection treatment)

IT Viral RNA

RL: BPR (Biological process); BIOL (Biological study); PROC (Process) (PEG-asparaginase, and combinations with other agents, for HIV infection treatment)

IT Polyoxalkylenes, biological studies

RL: BAC (Biological activity or effector, except adverse); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(asparaginase conjugates; PEG-asparaginase, and combinations with other agents, for HIV infection treatment)

IT Drug delivery systems

(prodrugs; PEG-asparaginase, and combinations with other agents, for HIV infection treatment)

EE Drug interactions

(mycogenic; PEG-asparaginase, and combinations with other agents, for HIV infection treatment)

ET 127-03-1, Hydroxyurea 3056-17-5, 64T 7481-39-2, Ddc 9015-68-3D, Asparaginase, PEG conjugates 25322-68-3D, PEG, asparaginase conjugates 30516-87-1, AZT 69655-05-6, Ddl 69639-83-4, Didox 74140-70-5, Paracetamol 95933-72-5, Amidox 95933-74-7, ***Trimidox*** 110313-27-8, Sulfonam 12318-82-1, 2-Chloro-9-(2-deoxy-2-fluoro-beta-D-erythrofuranylmethyl) 127142-14-7, BW 3489187 127779-20-8, Saquinavir 134306-02-4, NDL 101371 134674-17-4, 3TC 143621-35-6, 3-Aminocyclohexyl-2-carboxy-4-hydroxy-1-hydroxy-1-oxo-1,2,3,4-tetrahydro-1,2,3,4-tetrahydropyrimidin-2-one 344796-13-9, BILD 733 150378-17-9, Indinavir 150699-74-2, LY 295501 155213-67-5, Ritonavir 156130-57-3, LY 297702 157810-81-6, Crivarin 159999-64-7, Nelfinavir 159989-65-8, Viracept 160299-99-0, BILD 1263 161633-61-6, BILD 1357 161814-49-9, VIK-478 169238-52-2, BILD 1257 173943-67-9, 180100-89-6, TAS 106 180511-93-7, BILD 1351 211113-85-8 236391-65-4, BILD 1633 236391-66-5, GTI 2040 236391-67-6, GTI 2501 236392-18-0, Endinovere 236392-56-6, OCX 191 236392-75-9, PL 7

RL: BAC (Biological activity or effector, except adverse); THU

(Therapeutic use); BIOL (Biological study); USES (Uses)

(PEG-asparaginase, and combinations with other agents, for HIV infection treatment)

IT 56-84-8, L-Aspartic acid, biological studies 56-85-9, L-Glutamine,

biological studies 70-47-3, L-Asparagine, biological studies

RL: BOC (Biological occurrence); BPR (Biological process); BIOL

(Biological study); OCCU (Occurrence); PROC (Process)

(PEG-asparaginase, and combinations with other agents, for HIV infection treatment)

IT 9040-57-7, Ribonucleotide reductase 9068-38-6, Reverse transcriptase

144114-21-6, Retropepsin

RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL

(Biological study); PROC (Process)

(inhibitors; PEG-asparaginase, and combinations with other agents, for HIV infection treatment)

RE.CNT 8

RE

(1) Anil, T; JOURNAL OF ACQUIRED IMMUNE DEFICIENCY SYNDROMES AND HUMAN RETROVIROLOGY 1998, V17(4), PA31

(2) Gisselbrecht, C; AMERICAN JOURNAL OF MEDICINE 1993, V95(2), P188

MEDLINE

(3) Holle, L; ANNALS OF PHARMACOTHERAPY 1997, V31(5), P616 HCAPLUS

(4) Inst Nat Sante Rech Med; EP 0250335 A 1987

(5) Levien, T; HOSPITAL PHARMACY 1995, V30(1), P54

(6) Los Angeles Childrens Hospital; WO 9856410 A 1998

(7) Monfardini, S; CANCER TREATMENT REVIEWS 1994, V20(2), P149

(8) Tulpule, A; BLOOD 1998, V92(10), P240B

L3 ANSWER 6 OF 23 HCAPLUS COPYRIGHT 2000 ACS

AN 1999-113524 HCAPLUS

DN 130:177527

TI Therapeutic process for inhibiting NF-kappa.B

IN Elford, Howard L.

PA USA

SO PCT Int. Appl., 12 pp.

CODEN: PIXXD2

DT Patent

LA English

IC ICM A61K

CC 1-7 (Pharmacology)

FAN.CNT 1

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI WO 9906009	A2	19990211	WO 1998-US15715	19980729
WO 9906009	A3	19990902		

W: CA, JP

RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

PRAI US 1997-54230 19970730

OS MARPAT 130:177527

AB A therapeutic process is provided for the inhibition of NF-kappa.B in mammals in whose cells NF-kappa.B has been activated by an agency external to said cell.

ST nuclear factor kappaB inhibitor antiinflammatory

IT Animal virus

Antitumor agents

Arteriosclerosis

Chemotherapy

Diabetes mellitus

Oxidizing agents

Radiotherapy

Transplant (organ)

(NF-kappa.B activation by; therapeutic process for inhibiting NF-kappa.B)

IT Cytokines

RL: ADV (Adverse effect, including toxicity); BIOL (Biological study)

(NF-kappa.B activation by; therapeutic process for inhibiting NF-kappa.B)

IT NF-kappa.B

RL: ADV (Adverse effect, including toxicity); BPR (Biological process);

BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process)
(Inhibitors; therapeutic process for inhibiting NF- κ B) B)

II Anti-inflammatory drugs
Radical scavengers
(therapeutic process for inhibiting NF- κ B) B)

IT 143440-14-0, Protein kinase B
RL: ADP (Adverse effect, including toxicity); BIOL (Biological study)
(NF- κ B activation by therapeutic process for inhibiting NF- κ B) B)

IT 9040-57-7, Ribonucleotide reductase
RL: BSU (Biological study, unclassified); BIOL (Biological study)
(Inhibitors; therapeutic process for inhibiting NF- κ B) B)

II 95933-72-5, 95933-74-7, ***Trimidox***
RL: BAC (Biological activity or effector, except adverse); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(therapeutic process for inhibiting NF- κ B)

L3 ANSWER 7 OF 23 HCAPLUS COPYRIGHT 2000 ACS
AN 1998:727246 HCAPLUS
DN 130:90042

TI Characterization of the enzymic activity for biphasic competition by guanosine (1-(2,6-dichlorobenzylidene-amino)-3-hydroxyguanine) at .alpha.2-adrenoceptors. I. Description of an enzymic activity in spleen membranes

AU Uhlen, Staffan; Dambrova, Majja; Tiger, Gunnar; Oliver, Douglas W.; Wikberg, Jarl E. S.

CS Department of Pharmaceutical Biosciences, Division of Pharmacology, Uppsala University, Uppsala, Swed.

SO Biochem. Pharmacol. (1998), 56(9), 1111-1119
CODEN: BCPCA6; ISSN: 0006-2952

PB Elsevier Science Inc.

DT Journal

LA English

CC 1-2 (Pharmacology)

AB The mechanism for formation of high-affinity binding of 1-(2,6-dichlorobenzylidene-amino)-3-hydroxyguanine (guanosine) to .alpha.2-adrenoceptors was studied in particulate fractions from the rat spleen. The proportion of apparent high vs. low-affinity .alpha.2-adrenoceptor binding sites increased with increasing incubation time and was also augmented by Mg²⁺ ions. The formation of high-affinity guanosine binding seemed to be inhibited by a series of N-hydroxyguanine analogs to guanosine, as well as by a series of metabolic inhibitors that included allopurinol, 1-chloro-2,4-dinitrobenzene, 5,5'-dithiobis-(2-nitrobenzoic acid), cibacron blue, phenyl-p-benzoquinone, didox, and ***trimidox***. The formation of guanosine high-affinity binding was also inhibited in a time- and concn.-dependent fashion by preincubating the membranes with the LW03 N-hydroxyguanine analog of guanosine. Moreover, when the spleen membranes were extensively washed for 30 min with buffers at 25 degree., the guanosine high-affinity binding disappeared. However, when these washed membranes were supplemented with xanthine, the apparent affinity of guanosine increased four- to five-fold. Taken together, all data were compatible with the theory that the formation of high-affinity binding was dependent on the generation of a guanosine metabolite that showed an approx. 100-fold greater affinity for the .alpha.2-adrenoceptors than guanosine itself. Because the most potent blocker of the formation of high-affinity binding was allopurinol (apart from some N-hydroxyguanine analogs to guanosine) and since the activity could be restored with xanthine, a likely candidate responsible for the metabolic activation is xanthine oxidase.

ST guanosine metab spleen adrenoceptor binding; xanthine oxidase guanosine metab adrenoceptor binding

IT Cell membrane
Cerebral cortex
Drug metabolism
(metabolic activation in spleen and cerebral cortex membranes for guanosine binding to .alpha.2A-adrenoceptors)

IT Spleen
(metabolic activation in spleen membranes for guanosine binding to .alpha.2A-adrenoceptors)

IT .alpha.2-Adrenoceptors subtype A
RL: BPR (Biological process); BIOL (Biological study); PROC (Process)
(metabolic activation in spleen membranes for guanosine binding to .alpha.2A-adrenoceptors)

IT 9002-17-9, Xanthine oxidase
RL: BAC (Biological activity or effector, except adverse); BIOL (Biological study)
(metabolic activation in spleen membranes for guanosine binding to .alpha.2A-adrenoceptors)

IT 24047-25-4, Guanosine
RL: BAC (Biological activity or effector, except adverse); BPR (Biological process); BIOL (Biological study); PROC (Process)
(metabolic activation in spleen membranes for guanosine binding to .alpha.2A-adrenoceptors)

IT 7439-95-4, Magnesium, biological studies
RL: BOC (Biological occurrence); BIOL (Biological study); OCCU (Occurrence)
(metabolic activation in spleen membranes for guanosine binding to

.alpha.2A-adrenoceptors)

RECNT 23

RE

(1) Bergstrom, A; Acta Pharmacol Toxicol 1986, V59, P270 MEDLINE
(2) Carrington, H; Nature (London) 1951, V168, P1030
(3) Clement, B; Chem Res Toxicol 1998, V9, P662 HCAPLUS
(4) Crisafulli, D; Biochem Biophys Res Commun 1993, V191, P34 HCAPLUS
(5) Dambrova, M; Biochem Pharmacol 1998, V56, P1121 HCAPLUS
(6) De Lean, A; Mol Pharm 1983, V21, P3 HCAPLUS
(7) Desbail, P; Drug Res 1992, V42, P65 HCAPLUS
(8) Hotell, K; Ann Bio Chem (Paris) 1940, V31, P331 HCAPLUS
(9) Hui, M; Anticancer Res 1994, V24, P263 HCAPLUS
(10) Ledoux, F; Therapie 1981, V36, P187 MEDLINE
(11) Leri, F; Science (Washington DC) 1994, V266, P301 HCAPLUS
(12) Lowry, O. J; Biol Chem 1951, V193, P265
(13) Massey, V; J Biol Chem 1970, V245, P2837 HCAPLUS
(14) Uhlen, S; Br J Pharmacol 1991, V104, P657 HCAPLUS
(15) Uhlen, S; Br J Pharmacol 1992, V106, P986 HCAPLUS
(16) Uhlen, S; Eur J Pharmacol 1991, V202, P235 HCAPLUS
(17) Uhlen, S; Naunyn-Schmiedeberg Arch Pharmacol 1993, V347, P280 HCAPLUS
(18) Uhlen, S; Pharmacol Toxicol 1991, V69, P341 HCAPLUS
(19) Wang, P; J Med Chem 1990, V22, P608
(20) Ward, S; Br J Clin Pharmacol 1991, V31, P689 HCAPLUS
(21) Wikberg, J; Adrenoceptors 1995, P109 HCAPLUS
(22) Wikberg, J; Biochem Pharmacol 1998, V56, P1129 HCAPLUS
(23) Xia, Y; Pharmacol Toxicol 1993, V72, P40 HCAPLUS

L3 ANSWER 8 OF 23 HCAPLUS COPYRIGHT 2000 ACS
AN 1998:487631 HCAPLUS
DN 129:211378

TI Enhanced effects of Adriamycin by combination with a new ribonucleotide reductase inhibitor, ***trimidox***, in murine leukemia

AU Fritzer-Szekeres, Monika; Novotny, Ladislav; Romanova, Darina; Gobl, Rainer; Sedlak, Jan; Vachalkova, Anna; Rauko, Peter; Elford, Howard L.; Szekeres, Thomas

CS Clinical Institute for Medical and Chemical Laboratory Diagnostics, Vienna, A-1090, Austria

SO Life Sci. (1998), 63(7), 545-552
CODEN: LIFSAK; ISSN: 0024-3205

PB Elsevier Science Inc.

DT Journal

LA English

CC 1-6 (Pharmacology)

AB Ribonucleotide reductase is the rate limiting enzyme of de novo DNA synthesis; its activity is significantly increased in tumor cells related to the proliferation rate. Therefore the enzyme is considered to be an excellent target for cancer chemotherapy. In the present study we tested the in vitro and in vivo antitumor effects of a drug combination using ***trimidox*** (3,4,5-trihydroxybenzamidoxime), a novel inhibitor of ribonucleotide reductase with adriamycin, a widely used anticancer drug. This combination was selected because adriamycin generates free radicals being responsible for cardiotoxic side effects, ***trimidox*** has been shown to be a good free radical scavenger. The in vitro cytotoxic effect of the drug combination was examd. in L1210 mouse leukemia cells employing a MTT chemosensitivity assay. Incubation of these cells with adriamycin and ***trimidox*** together yielded less than additive cytotoxic effects compared to either drug alone. These effects were not caused by the involvement of p-glycoprotein mediated drug efflux. However, when the effect of ***trimidox*** and adriamycin in combination was examd. in L1210 leukemia bearing mice antitumor effects of adriamycin could be enhanced by the presence of ***trimidox***. Our data indicate, that the in vivo combination of adriamycin together with ***trimidox*** might be beneficial for the treatment of malignancies.

ST ***trimidox*** Adriamycin leukemia inhibition

IT Drug transport
(P-glycoprotein-mediated; enhanced effects of Adriamycin by combination with a new ribonucleotide reductase inhibitor, ***trimidox***, in murine leukemia)

IT Leukemia inhibitors
Synergistic drug interactions
(enhanced effects of Adriamycin by combination with a new ribonucleotide reductase inhibitor, ***trimidox***, in murine leukemia)

IT 25316-40-9, Adriamycin 95933-74-7, ***Trimidox***
RL: BAC (Biological activity or effector, except adverse); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(enhanced effects of Adriamycin by combination with a new ribonucleotide reductase inhibitor, ***trimidox***, in murine leukemia)

IT 9040-57-7, Ribonucleotide reductase
RL: BSU (Biological study, unclassified); BIOL (Biological study)
(enhanced effects of Adriamycin by combination with a new ribonucleotide reductase inhibitor, ***trimidox***, in murine leukemia)

L3 ANSWER 9 OF 23 HCAPLUS COPYRIGHT 2000 ACS
AN 1998:429154 HCAPLUS
DN 129:170304

TI ***Trimidox*** -mediated morphological changes during erythroid differentiation is associated with the stimulation of hemoglobin and

F-cell production in human K562 cells

AU Iyama, Etsu W.; Adeniyi, Samuel E.; Elford, Howard L.; Foulds, Hugo; Turner, Ernest A.

CS Comprehensive Sickle Cell Center, Nashville, TN, 37203, USA

SO Biochem Biophys Res Commun. (1998) 247(3): 739-704

CODEN: BBMLCA; ISSN: 0006-291X

PR Academic Press

DT Journal

LA English

CC 1-6 (Pharmacology)

AB ***Trimidox*** (7,4,5-trihydroxybenzohydroxamide) has been shown to reduce the activity of ribonucleotide reductase with accompanying growth inhibition and differentiation of mammalian cells. Hydroxyurea (HU) is the only ribonucleotide reductase inhibitor in clin. use for the treatment and management of sickle cell anemia, since this compd. increases fetal Hb (Hb F) prodn.: a potent inhibitor of sickle Hb (Hb,SS) polymn. However, the main limitations of HU is its lack of potency, myelosuppression and short half life. These studies investigated the effects of ***trimidox*** on the induction of Hb and F-cells prodn. in K562 erythroleukemia cells. Our study reveals that ***trimidox*** exhibits concn. dependent inhibitory effect on K562 cells with increase in benzidine pos. normoblasts and F-cells prodn. as well as morphol. changes typical of erythroid differentiation. These findings provide the first evidence that the growth inhibitory differentiation of cells induced by ***trimidox*** enhance Hb and F-cells prodn. (c) 1998 Academic Press.

ST ***trimidox*** fetal Hb ribonucleotide reductase; sickle Hb

IT Erythrocyte (differentiation; ***trimidox*** -mediated morphol. changes during erythroid differentiation is assoc. with the stimulation of Hb and F-cell prodn. in human K562 cells)

IT Sickle cell anemia (***trimidox*** -mediated morphol. changes during erythroid differentiation is assoc. with the stimulation of Hb and F-cell prodn. in human K562 cells)

IT 95933-74-7, ***Trimidox***

RL: ADV (Adverse effect, including toxicity); BAC (Biological activity or effector, except adverse); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(***trimidox*** -mediated morphol. changes during erythroid differentiation is assoc. with the stimulation of Hb and F-cell prodn. in human K562 cells)

IT 9034-63-3, Hemoglobin F 9040-57-7, Ribonucleotide reductase

RL: BPR (Biological process); BIOL (Biological study); PROC (Process)

(***trimidox*** -mediated morphol. changes during erythroid differentiation is assoc. with the stimulation of Hb and F-cell prodn. in human K562 cells)

L3 ANSWER 10 OF 23 HCAPLUS COPYRIGHT 2000 ACS

AN 1998:415518 HCAPLUS

DN 129:156586

TI Interaction of gallium nitrate with other inhibitors of ribonucleotide reductase: effects on the proliferation of human leukemic cells

AU Myette, Michael S.; Elford, Howard L.; Chitambar, Christopher R.

CS Division of Hematology/Oncology, Medical College of Wisconsin, Milwaukee, WI, 53226, USA

SO Cancer Lett. (Shannon, Irel.) (1998), 129(2), 199-204

CODEN: CALEDQ; ISSN: 0304-3835

PB Elsevier Science Ireland Ltd.

DT Journal

LA English

CC 1-6 (Pharmacology)

AB Ribonucleotide reductase, a key enzyme in deoxyribonucleotide synthesis, is an important target for cancer chemotherapy. Drugs that inhibit its individual components may act synergistically to block DNA synthesis. Prior work has established that gallium inhibits the R2 subunit of ribonucleotide reductase. We show that gallium acts synergistically with the ribonucleotide reductase inhibitors gemcitabine and hydroxyurea to inhibit the proliferation of CCRF-CEM cells. In contrast, combinations of gallium with the ribonucleotide reductase inhibitors amidox, didox, or ***trimidox*** produced antagonistic effects on cell growth. Spectroscopy anal. revealed that as a result of their metal-binding properties, amidox, didox and ***trimidox*** formed complexes with gallium, thus negating potential synergistic actions. Our results have important implications in the design of clin. trials using these ribonucleotide reductase inhibitors in combination.

ST ribonucleotide reductase inhibitor gallium leukemia

IT DNA formation

Leukemia inhibitors

Synergistic drug interactions

(interaction of gallium nitrate with other inhibitors of ribonucleotide reductase and effects on proliferation of human leukemic cells)

IT 127-07-1, Hydroxyurea 13494-90-1, Gallium nitrate 69839-83-4, Didox 95058-81-4, Gemcitabine 95933-72-5, Amidox 95933-74-7, ***Trimidox***

RL: BAC (Biological activity or effector, except adverse); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(interaction of gallium nitrate with other inhibitors of ribonucleotide reductase and effects on proliferation of human leukemic cells)

IT 9040-57-7, Ribonucleotide reductase

RL: BSU (Biological study, unclassified); BIOL (Biological study)

(interaction of gallium nitrate with other inhibitors of ribonucleotide reductase and effects on proliferation of human leukemic cells)

L3 ANSWER 11 OF 23 HCAPLUS COPYRIGHT 2000 ACS

AN 1998:69701 HCAPLUS

DN 128:169455

TI DNA-protective activity of new ribonucleotide reductase inhibitors

AU Raulo, Peter; Romanova, Darina; Machkova, Eva; Mochalov, Kvetoslav; Novotny, Ladislav; Elford, Howard L.; Szekeres, Thomas

CS Department of Experimental Therapy, Cancer Research Institute, Slovak Academy of Sciences, Bratislava, SK-81232, Slovakia

SO Anticancer Res. (1997), 17(SA), 3437-3440

CODEN: ANTRD4; ISSN: 0250-7095

PB Anticancer Research

DT Journal

LA English

CC 1-12 (Pharmacology)

AB The DNA-protective activity of hydroxyurea (HU) and novel ribonucleotide reductase (RR) inhibitors amidox (AX), didox (DX) and ***trimidox*** (TX) was examd. using hydrogen peroxide as the DNA-damaging agent. The exposure of superspiralized plasmid DNA mols. (pBR 322) to H2O2 under precisely defined in vitro conditions initiates a change in DNA topol. (DNA form I relaxes to DNA form II). This electrophoretically monitored change in the plasmid DNA topol. is related to the induction of ss-DNA breaks and corresponds with DNA exposition to free radicals. The inhibition of DNA relaxation (the prevention of DNA damage induced by hydrogen peroxide) depended on the free radical scavenging capacity of the drugs investigated. HU exerted DNA protective activity at a concn. of 4 mM, AX at concn. of 1 .mu.M, TX at a concn. of 5 .mu.M and DX at a concn. of 25 .mu.M (the free radical scavenging activity increases from HU to AX in following manner: HU < mcht. DX < TX < AX). It can be concluded that the new synthetic RR-inhibitor AX which is being investigated at the preclin. level as a potential anti-cancer drug possess the highest capacity for scavenging of free radicals.

ST DNA protection ribonucleotide reductase inhibitor; radical scavenging ribonucleotide reductase inhibitor DNA

IT DNA damage

Radical scavengers

(DNA-protective activity of new ribonucleotide reductase inhibitors and hydroxyurea in relation to radical scavenging capacity)

IT 7722-84-1, Hydrogen peroxide, biological studies

RL: ADV (Adverse effect, including toxicity); BPR (Biological process); BIOL (Biological study); PROC (Process)

(DNA-protective activity of new ribonucleotide reductase inhibitors and hydroxyurea in relation to radical scavenging capacity)

IT 127-07-1, Hydroxyurea 69839-83-4, Didox 95933-72-5, Amidox

RL: BAC (Biological activity or effector, except adverse); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(DNA-protective activity of new ribonucleotide reductase inhibitors and hydroxyurea in relation to radical scavenging capacity)

IT 9040-57-7, Ribonucleotide reductase

RL: BSU (Biological study, unclassified); BIOL (Biological study)

(inhibitors; DNA-protective activity of new ribonucleotide reductase inhibitors and hydroxyurea in relation to radical scavenging capacity)

L3 ANSWER 12 OF 23 HCAPLUS COPYRIGHT 2000 ACS

AN 1998:26676 HCAPLUS

DN 128:136198

TI Enhanced effects of adriamycin by combination with a new ribonucleotide reductase inhibitor, ***trimidox***, in murine leukemia

AU Novotny, L.; Romanova, D.; Gobl, R.; Sedlak, J.; Vachalkova, A.; Rauko, P.; Fritzer-Szekeres, M.; Elford, H. L.; Szekeres, T.

CS Cancer Research Inst., SAS, Bratislava, SK-812 32, Slovakia

SO Haematol. Blood Transfus. (1998), 39(Acute Leukemias VII), 556-561

CODEN: HBTRDV; ISSN: 0171-7111

PB Springer-Verlag

DT Journal

LA English

CC 1-6 (Pharmacology)

AB Ribonucleotide reductase is the rate limiting enzyme of de novo DNA synthesis; its activity is significantly increased in tumor cells related to the proliferation rate of the tumor cell. Therefore the enzyme is considered to be an excellent target for cancer chemotherapy. In the present study we tested the in vitro and in vivo antitumor effects of a drug combination using ***trimidox*** (3,4,5-trihydroxybenzohydroxamide), a novel inhibitor of ribonucleotide reductase with adriamycin, a widely used anticancer drug. This combination was selected because adriamycin generates free radicals, which are responsible for cardiotoxic side effects of adriamycin treatment, and because ***trimidox*** has been shown to be a good free radical scavenger. The in vitro cytotoxic effect of the drug combination was examd. in L 1210 mouse leukemia cells employing an MTT chemo-sensitivity assay. Simultaneous in vitro incubation of these cells yielded antagonistic cytotoxic effects compared to either drug alone. These effects were not caused by the involvement of p-glycoprotein mediated drug efflux. However, when the effect of ***trimidox*** and adriamycin in combination was examd. in L 1210 leukemia bearing mice, antitumor effects of adriamycin could be enhanced by the presence of ***trimidox***. Animals were treated on day two after tumor cell injection with 5 mg/kg

adriamycin and received 250 mg/kg ***trimidox*** on days 2,3 and 6. Mice treated with adriamycin or ***trimidox*** alone yielded a 41 and 34% increase in life span, resp. However, animals which were treated with both drugs, showed a 65% increase of their life span. Our data indicate, that in vitro results of drug combinations should be interpreted with extreme caution and suggest that the in vivo combination of adriamycin together with ***trimidox*** might be beneficial for the treatment of malignancies.

ST Adriamycin treatment ***trimidox*** P glycoprotein

IT Drug interactions

Leukemia inhibitors

Adriamycin antileukemic effects enhanced by ribonucleotide reductase inhibitor ***trimidox***)

II P-glycoproteins

RL: BSU (Biological study, unclassified); BIOL (Biological study) (adriamycin antileukemic effects enhancement by ribonucleotide reductase inhibitor ***trimidox***)

IT 25316-40-9, Adriamycin 95933-74-7, ***Trimidox***

RL: BAC (Biological activity or effector, except adverse); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (adriamycin antileukemic effects enhancement by ribonucleotide reductase inhibitor ***trimidox***)

L3 ANSWER 13 OF 23 HCAPLUS COPYRIGHT 2000 ACS

AN 1997:795554 HCAPLUS

DN 128:123476

TI Effective use of ribonucleotide reductase inhibitors (didox and ***trimidox***) alone or in combination with didanosine (ddI) to suppress disease progression and increase survival in murine acquired immunodeficiency syndrome (MAIDS)

AU Mayhew, Christopher; Oakley, Oliver; Piper, James; Hughes, Nedda K.; Phillips, Jonathan; Birch, Nicholas J.; Elford, Howard L.; Gallicchio, Vincent S.

CS Laboratory of Experimental Immunohematopoiesis and Developmental Therapeutics, Departments of Clinical Sciences and Internal Medicine, Chandler Medical Center, University of Kentucky, Lexington, KY, 40536, USA

SO Cell. Mol. Biol. (Paris) (1997), 43(7), 1019-1029

CODEN: CMOBEF; ISSN: 0145-5680

PB C.M.B. Association

DT Journal

LA English

CC 1-5 (Pharmacology)

AB Ribonucleotide reductase inhibitors (RRIs) have been recently shown to inhibit retroviral replication. We examd. a new series of RRIs, 3,4-dihydroxybenzohydroxamic acid (Didox) and 3,4,5-trihydroxybenzohydroxamidoxime (***Trimidox***) for their ability to alter disease progression in murine acquired immunodeficiency syndrome (MAIDS), both alone and in combination with 2',3'-dideoxyinosine (ddI). MAIDS disease was induced by inoculation of female C57BL/6 mice with the LP-BMS murine leukemia virus (MuLV) and disease progression characterized by extensive peripheral lymphadenopathy and splenomegaly. Efficacy of treatment with these drugs was based upon their ability to influence survival and disease pathophysiol. by monitoring the development of splenomegaly. Toxicity was detd. by changes in body wt., total peripheral white blood cell count and hematocrit. Didox or ***trimidox*** monotherapy was assocd. with increased survival and decreased disease pathophysiol., with no apparent toxicity. Combined with ddI, their ability to reduce development of viral induced splenomegaly was enhanced compared to ***trimidox***, didox or ddI alone. These results demonstrate RRIs have potent activity in reversing the disease manifestations characteristic of MAIDS. Further studies are warranted to det. human clin. efficacy.

ST ribonucleotide reductase inhibitor didanosine murine AIDS; antiviral didox ***trimidox*** didanosine AIDS HIV1

IT AIDS (disease)

Antiviral agents

Drug interactions

Human immunodeficiency virus 1

(ribonucleotide reductase inhibitors (didox and ***trimidox***) alone or in combination with didanosine: suppression of MAIDS)

IT 9040-57-7, Ribonucleotide reductase

RL: BSU (Biological study, unclassified); BIOL (Biological study) (inhibitors; ribonucleotide reductase inhibitors (didox and ***trimidox***) alone or in combination with didanosine: suppression of MAIDS)

IT 69655-05-6, Didanosine 69839-83-4, Didox 95933-74-7, ***Trimidox***

RL: BAC (Biological activity or effector, except adverse); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (ribonucleotide reductase inhibitors (didox and ***trimidox***) alone or in combination with didanosine: suppression of MAIDS)

L3 ANSWER 14 OF 23 HCAPLUS COPYRIGHT 2000 ACS

AN 1997:567538 HCAPLUS

DN 127:243220

TI Selective inhibition of I.kappa.B.alpha. phosphorylation and HIV-1 LTR-directed gene expression by novel antioxidant compounds

AU Lee, Raymond; Beauparlant, Pierre; Elford, Howard; Ponka, Premysl; Hiscott, John

CS Lady Davis Institute for Medical Research, McGill University, Montreal, PQ, H3T 1E2, Can.

SO Virology (1997), 234(2), 377-390

CODEN: VIRLAX; ISSN: 0042-6822

PB Academic

DT Journal

LA English

CC 1-12 (Pharmacology)

AB Oxidative stress activates the NF-kappa.B/NFkB transcription factors which are involved in the activation of numerous immunoregulatory genes and the human immunodeficiency virus type 1 (HIV-1) long terminal repeat (LTR). In the present study, we examined the effects of established and novel compounds, including antioxidants, ribonucleotide reductase inhibitors, and iron chelators on NF-kappa.B activation and HIV LTR-mediated gene expression induced by TNF-alpha, N-Acetylcysteine (NAC), pyrrolidinecarboxamide (PDC), and ***Trimidox*** (TD) at various concentrations. Pretreatment of cells with these compounds prior to stimulation prevented I.kappa.B.alpha. degradn. Phosphorylation of I.kappa.B.alpha., a prerequisite for its signal-induced degradn., was abrogated in these cells, indicating that oxidative stress is an essential step in the NF-kappa.B activation pathway. On the other hand, iron chelators desferrioxamine, pyridoxal isonicotinoyl hydrazone (PIH), and salicylaldehyde isonicotinoyl hydrazone (SIH) showed no inhibition of TNF-alpha-induced NF-kappa.B DNA-binding activity. Synergistic induction of HIV-1 LTR-mediated gene expression by TNF-alpha and the HIV-1 transactivator Tat in Jurkat cells was significantly suppressed in the presence of NAC and TD, but not PDC. The inhibition of NAC and TD on LTR-directed gene expression was diminished when NF-kappa.B-binding sites in the LTR were deleted, indicating that these compounds affected the NF-kappa.B component of the synergism. Iron chelators PIH and SIH also showed some inhibitory effect on LTR-mediated gene activation, presumably through an NF-kappa.B-independent mechanism. These expts. demonstrate that TD, at concn. 50 times lower than the effective concn. of NAC, potently inhibits NF-kappa.B activity and suppresses HIV LTR expression.

ST antioxidant NFkB HIV1 gene

IT Antioxidants (pharmaceutical)

Human immunodeficiency virus 1

(inhibition of I.kappa.B.alpha. phosphorylation and HIV-1 LTR-directed gene expression by antioxidants)

IT Tumor necrosis factor .alpha.

RL: BAC (Biological activity or effector, except adverse); BIOL (Biological study)

(inhibition of I.kappa.B.alpha. phosphorylation and HIV-1 LTR-directed gene expression by antioxidants)

IT LTR (long terminal repeat)

RL: BPR (Biological process); BIOL (Biological study); PROC (Process) (inhibition of I.kappa.B.alpha. phosphorylation and HIV-1 LTR-directed gene expression by antioxidants)

IT NF-kappa.B

RL: BPR (Biological process); BIOL (Biological study); PROC (Process) (inhibition of I.kappa.B.alpha. phosphorylation and HIV-1 LTR-directed gene expression by antioxidants)

IT Chelating agents

(iron; inhibition of I.kappa.B.alpha. phosphorylation and HIV-1 LTR-directed gene expression by antioxidants)

IT 70-51-9, Desferrioxamine 495-84-1, Salicylaldehyde isonicotinoyl hydrazone 616-91-1, N-Acetylcysteine 737-86-0, Pyridoxal isonicotinoyl hydrazone 25769-03-3, 1-Pyrrolidinecarboxylic acid 69839-83-4, Didox 95933-72-5, Amidox 95933-74-7, ***Trimidox***

RL: BAC (Biological activity or effector, except adverse); BIOL (Biological study)

(inhibition of I.kappa.B.alpha. phosphorylation and HIV-1 LTR-directed gene expression by antioxidants)

L3 ANSWER 15 OF 23 HCAPLUS COPYRIGHT 2000 ACS

AN 1997:529356 HCAPLUS

DN 127:130355

TI The effect of new combinations of antimetabolites and ***trimidox*** on cancer cells

AU Romanova, D.; Raslova, H.; Plaschke, K.; Novotny, L.; Fritzer, M.

CS Ustav experimentalnej onkologie, Bratislava, 812 32, Slovakia

SO Farm. Obz. (1995), 64(7-8), 180-187

CODEN: FAOBAS; ISSN: 0014-8172

PB Zdravotnicke Vydavatelstvo HERBA

DT Journal; General Review

LA Slovak

CC 1-0 (Pharmacology)

AB A review with 22 refs. The effects of ***trimidox***, a new inhibitor of ribonucleotide reductase, used in combination with antimetabolites arabinosylcytosine (ara-C) and gemcitabine (difluorodeoxycytidine) used in anticancer chemotherapy were studied in vitro cultures of human colon cancer HT-29 cells. The effects ***trimidox*** were compared with the effects of thiazofurine combined with hypoxanthine or allopurinol. The cytostatic effects were also evaluated in human leukemic cells HL-60. The levels of ribonucleoside and deoxyribonucleoside triphosphates and cell cycle responses were detd. The mechanisms of ***trimidox*** action, biochem. pathways, anticancer activity, synergism, and cytotoxicity are discussed.

ST review ***trimidox*** antitumor combination araC gemcitabine

IT Antitumor agents

Drug interactions

→ Aug. 4th

- HL-60 cell
HT-29 cell
(antitumor effect of ***trimidox*** in combination of antineoplastic agents in cancer cells)
- IT 63-94-0, Hypoxanthine 127-07-1, Hydroxyurea 147-94-1, Ara-C 113-30-0, Allopurinol 60034-10-8, Tracofurin 69839-83-4, Didox 95933-72-5, Gemcitabine 95933-74-7, ***Trimidox***
RL: BAC (Biological activity or effector, except adverse); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(antitumor effect of ***trimidox*** in combination of antineoplastic agents in cancer cells)
- L3 ANSWER 16 OF 23 HCAPLUS COPYRIGHT 2000 ACS
AN 1997:99918 HCAPLUS
DN 127:13317
TI Genotoxic properties of the newly synthesized antineoplastic agents amidox, didox, and ***trimidox***
AU Mladkova, E.; Macakova, K.; Podstavkova, S.; Vlcek, D.
CS Department Genetics, Faculty Sciences, Bratislava, 84215, Slovakia
SO Pharmazie (1997), 52(7), 540-544
CODEN: PHARAT; ISSN: 0031-7144
PB Govi-Verlag Pharmazeutischer Verlag
DT Journal
LA English
CC 1-6 (Pharmacology)
Section cross-reference(s): 4
AB Toxic and genotoxic effects of 3 polyhydroxy-substituted benzohydroxamates (amidox, didox, and ***trimidox***), having antineoplastic activities by the mechanism of the ribonucleotide reductase activity inhibition, were evaluated by reverse mutation assay on Salmonella typhimurium strains TA97, TA98, TA100, TA102. While amidox did not exert any toxic effect, didox, and ***trimidox*** were toxic. The toxicity of the test chems. was dependent on the structure of their mol. and the repair capacity of the test strains. ***Trimidox*** exhibited the highest toxicity, and it was proved as a direct-acting frameshift mutagen. Its mutagenic effect was increased after a metabolic activation. Amidox and didox can be classified as frameshift promutagens.
ST antineoplastic agent amidox didox ***trimidox*** genotoxicity
IT Frameshift mutation
(genotoxicity of antineoplastic agent ***trimidox*** caused by)
IT Antitumor agents
(genotoxicity of antineoplastic agents amidox, didox, and ***trimidox***)
IT Genotoxicity
Toxicity
(of antineoplastic agents amidox, didox, and ***trimidox***)
IT 69839-83-4, Didox 95933-72-5, Amidox 95933-74-7, ***Trimidox***
RL: ADV (Adverse effect, including toxicity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(genotoxicity of antineoplastic agents)
IT 9047-64-7
RL: BPR (Biological process); BIOL (Biological study); PROC (Process)
(genotoxicity of antineoplastic agents amidox, didox, and ***trimidox*** caused by inhibition of)
- L3 ANSWER 17 OF 23 HCAPLUS COPYRIGHT 2000 ACS
AN 1997:32717 HCAPLUS
DN 127:39615
TI The new inhibitors of ribonucleotide reductase. Comparison of some physicochemical properties
AU Romanova, Darina; Vachalkova, Anna; Szekeres, Thomas; Elford, Howard L.; Novotny, Ladislav
CS Cancer Res. Inst. Slovak Academy Sci., Bratislava, SK-81232, Slovakia
SO J. Pharm. Biomed. Anal. (1997), 15(7), 951-956
CODEN: JPBAAD; ISSN: 0731-7085
PB Elsevier
DT Journal
LA English
CC 63-5 (Pharmaceuticals)
Section cross-reference(s): 22
AB Amidox (AX), didox (DX) and ***trimidox*** (TX), compds. synthesized as new ribonucleotide reductase inhibitors, have been investigated by UV spectrophotometry, polarog. HPLC. The expts. were performed at various pH values. The changes in UV absorption of the compds. studied were recorded and it was demonstrated that these changes are related to the pH and to structural features of the investigated mols. Only amidox and ***trimidox*** were reduced during polarog. expts. in Britton-Robinson buffer. The redn. of both compds. proceeded in 2 1-electron steps in acid solns. One 2-electron diffuse irreversible wave was obsd. at basic pH values. The values of the half-wave potential became more neg. with increasing pH values. HPLC assay also showed changes in the retention of compds. investigated, particularly when the pH of the mobile phase was close to the disson. const. of the particular drug. The changes of physicochem. properties detected by the methods are related to different chem. structures (the most significant changes were obsd. in alk. pH).
ST ribonucleotide reductase inhibitor physicochem property; polarog ribonucleotide reductase inhibitor; UV spectrometry ribonucleotide reductase inhibitor
IT Electrochemical reduction
UV and visible spectroscopy
(physicochem. properties of ribonucleotide reductase inhibitors)
IT 6040-52-7, Ribonucleotide reductase
RL: BPR (Biological process); BIOL (Biological study)
(physicochem. properties of ribonucleotide reductase inhibitors)
IT 69839-83-4, Didox 95933-72-5, Amidox 95933-74-7, ***Trimidox***
RL: BPR (Biological process); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(physicochem. properties of ribonucleotide reductase inhibitors)
- L3 ANSWER 18 OF 23 HCAPLUS COPYRIGHT 2000 ACS
AN 1995:1007445 HCAPLUS
DN 124:15813
TI Iron binding capacity of ***trimidox*** (3,4,5-trihydroxybenzamidoxime), a new inhibitor of the enzyme ribonucleotide reductase
AU Szekeres, Thomas; Vielnascher, Elisabeth; Novotny, Ladislav; Vachalkova, Anna; Fritzer, Monika; Findenig, Gabriele; Goebel, Rainer; Elford, Howard L.; Goldenberg, Hans
CS Inst. Medizinische Chemie, Univ. Wien, Vienna, Austria
SO Eur. J. Clin. Chem. Clin. Biochem. (1995), 33(11), 785-9
CODEN: EJCBEQ; ISSN: 0939-4974
DT Journal
LA English
CC 1-6 (Pharmacology)
AB Ribonucleotide reductase is the rate limiting enzyme of deoxynucleoside triphosphate synthesis and is considered to be an excellent target of cancer chemotherapy. ***Trimidox***, a newly synthesized compd., inhibits this enzyme and has in vitro and in vivo antitumor activity. As ***trimidox*** was able to upregulate the expression of the transferrin receptor in HL-60 human promyelocytic leukemia cells, the authors have now investigated the capability of ***trimidox*** to interfere with iron metab. The authors show by photometric and polarog. methods that ***trimidox*** is able to form an iron complex. However, its cytotoxic action cannot be circumvented by addn. of iron-satd. transferrin or iron-ammonium citrate, indicating that the iron complexing capacity is not responsible for the mechanism of action of this compd. When HL-60, K562 or L1210 leukemia cells were incubated with the ***trimidox***-iron complex itself, the authors could observe increases of the 50% growth inhibitory capacity of the complex in comparison with ***trimidox*** alone. The authors conclude that ***trimidox*** is able to form an iron complex, but in contrast to other agents, the anticancer activity cannot be contributed to this effect alone. Further studies will have to elucidate the mol. mechanism of action of this new and promising anticancer agent.
ST iron ***trimidox*** complex ribonucleotide reductase antitumor
IT Neoplasm inhibitors
(iron binding capacity of ***trimidox*** (3,4,5-trihydroxybenzamidoxime), a new inhibitor of the enzyme ribonucleotide reductase)
IT 9068-66-0, Ribonucleotide reductase
RL: BPR (Biological process); BIOL (Biological study); PROC (Process)
(inhibitor; iron binding capacity of ***trimidox*** (3,4,5-trihydroxybenzamidoxime), a new inhibitor of the enzyme ribonucleotide reductase)
IT 1185-57-5D, Ferric ammonium citrate, ***trimidox*** complex
RL: BAC (Biological activity or effector, except adverse); BIOL (Biological study)
(iron binding capacity of ***trimidox*** (3,4,5-trihydroxybenzamidoxime), a new inhibitor of the enzyme ribonucleotide reductase)
IT 95933-74-7, ***Trimidox***
RL: BAC (Biological activity or effector, except adverse); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(iron binding capacity of ***trimidox*** (3,4,5-trihydroxybenzamidoxime), a new inhibitor of the enzyme ribonucleotide reductase)
IT 7439-89-6, Iron, biological studies
RL: BPR (Biological process); BIOL (Biological study); PROC (Process)
(iron binding capacity of ***trimidox*** (3,4,5-trihydroxybenzamidoxime), a new inhibitor of the enzyme ribonucleotide reductase)
- L3 ANSWER 19 OF 23 HCAPLUS COPYRIGHT 2000 ACS
AN 1995:982076 HCAPLUS
DN 124:134431
TI Ribonucleotide reductase as target for enzyme-directed chemotherapy. Effects of ***trimidox*** (3,4,5-trihydroxybenzohydroxamidoxime), a new inhibitor of ribonucleotide reductase
AU Findenig, G.; Vielnascher, E.; Goebel, R.; Fritzer-Szekeres, M.; Szekeres, T.
CS Inst. Med. Chem., Univ. Wien, Vienna, A-1090, Austria
SO Wien. Klin. Wochenschr. (1995), 107(22), 694-7
CODEN: WKWOAO; ISSN: 0043-5325
DT Journal; General Review
LA German
CC 1-0 (Pharmacology)
Section cross-reference(s): 7
AB A review with 28 refs. describing the biochem., morphol., and cytotoxic effects of ***trimidox*** and other polyhydroxy-substituted benzohydroxamate derivs. on leukemia cell lines. Selection criteria,

- effects, and combinations used to enzyme-targeted chemotherapy are described for these ribonucleotide reductase inhibitors.
- S1 Tiazofurin benzohydroxamido cancer chemotherapy. ***Trimidox*** blockaden cytotoxic effect leukemia review, ribonucleotide reductase inhibitor ***Trimidox*** overview
- I1 Neoplasm inhibitors (ribonucleotide reductase as target for enzyme-directed chemotherapy)
- I1 95933-74-7, ***Trimidox***
 RL: BAC (Biological activity or effector, except adverse); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
 (ribonucleotide reductase as target for enzyme-directed chemotherapy)
- I1 9047-64-7, Ribonucleotide reductase
 RL: BSU (Biological study, unclassified); BIOL (Biological study)
 (ribonucleotide reductase as target for enzyme-directed chemotherapy)
- L3 ANSWER 20 OF 23 HCAPLUS COPYRIGHT 2000 ACS
 AN 1995:270272 HCAPLUS
 DN 122:45904
 TI Synergistic growth inhibitory and differentiating effects of ***Trimidox*** and tiazofurin in human promyelocytic leukemia HL-60 cells
 AU Szekeres, Thomas; Fritzer, Monika; Strobl, Herbert; Gharehbaghi, Kamran; Findenig, Gabriele; Elford, Howard L.; Lhotka, Christian; Schoen, Hans J.; Jayaram, Hiremagalur N.
 CS Inst. Med. Chem., Univ. Vienna Med. Sch., Vienna, Austria
 SO Blood (1994), 84(12), 4316-21
 CODEN: BLOODAW; ISSN: 0006-4971
 DT Journal
 LA English
 CC 1-6 (Pharmacology)
 AB Increased ribonucleotide reductase (RR) activity has been linked with malignant transformation and tumor cell growth. Therefore, this enzyme is considered to be an excellent target for cancer chemotherapy. The authors have examd. the effects of a newly patented RR inhibitor, ***Trimidox*** (3,4,5-trihydroxybenzohydroxamidoxime). ***Trimidox*** inhibited the growth of human promyelocytic leukemia HL-60 cells with an IC50 of 35 .mu.mol/L. Incubation of HL-60 cells with 50 .mu.mol/L ***Trimidox*** for 24 h decreased deoxyguanosine triphosphate (dGTP) and deoxycytidine triphosphate (dCTP) pools to 24% and 39% of control values, resp. Incubation of HL-60 cells with 20 to 80 .mu.mol/L ***Trimidox*** even up to a period of 4 days did not alter the distribution of cells in different phases of cell cycle. Sequential incubation of HL-60 cells with ***Trimidox*** (25 .mu.mol/L) for 24 h and then with 10 .mu.mol/L tiazofurin (an inhibitor of inosine monophosphate dehydrogenase) for 4 days produced synergistic growth inhibitory activity, and the cell no. decreased to 16% of untreated controls. When differentiation-linked cell surface marker expressions were detd. in cells treated with ***Trimidox*** and tiazofurin, a significantly increased fluorescence intensity was obsd. for the CD 11b (2.9-fold), CD 33 (1.9-fold), and HLA-D cell surface antigens. Expression of the transferrin receptor (CD71) increased 7.3-fold in cells treated with both agents, compared with untreated controls. The results suggest that ***Trimidox*** in combination with tiazofurin might be useful in the treatment of leukemia.
- ST promyelocytic leukemia inhibitor ***Trimidox*** tiazofurin synergism
 IT Transferrin receptors
 RL: BSU (Biological study, unclassified); BIOL (Biological study)
 (synergistic growth inhibitory and differentiating effects of ***Trimidox*** and tiazofurin in human promyelocytic leukemia HL-60 cells)
 IT Neoplasm inhibitors (promyelocytic leukemia, synergistic growth inhibitory and differentiating effects of ***Trimidox*** and tiazofurin in human promyelocytic leukemia HL-60 cells)
 IT Drug interactions (synergistic, synergistic growth inhibitory and differentiating effects of ***Trimidox*** and tiazofurin in human promyelocytic leukemia HL-60 cells)
 IT Receptors
 RL: BSU (Biological study, unclassified); BIOL (Biological study)
 (transferrin, synergistic growth inhibitory and differentiating effects of ***Trimidox*** and tiazofurin in human promyelocytic leukemia HL-60 cells)
 IT 60084-10-8, Tiazofurin 95933-74-7, ***Trimidox***
 RL: BAC (Biological activity or effector, except adverse); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
 (synergistic growth inhibitory and differentiating effects of ***Trimidox*** and tiazofurin in human promyelocytic leukemia HL-60 cells)
 IT 9040-57-7, Ribonucleotide reductase
 RL: BSU (Biological study, unclassified); BIOL (Biological study)
 (synergistic growth inhibitory and differentiating effects of ***Trimidox*** and tiazofurin in human promyelocytic leukemia HL-60 cells)
- L3 ANSWER 21 OF 23 HCAPLUS COPYRIGHT 2000 ACS
 AN 1995:79317 HCAPLUS
 DN 123:102138
 TI Synergistic cytotoxic effects of chemotherapeutic drugs on colon tumor cells by simultaneous inhibition of de novo and salvage metabolic pathways
 AU Szekeres, T.; Fritzer, M.; Schoen, H. J.; Findenig, G.; Lhotka, C.
- CS Inst. Med. Chem., Univ. Wien, Vienna, A-1090, Austria
 SO Wien. Klin. Wochenschr. (1994), 106(14), 459-63
 CODEN: WKWDAO; ISSN: 0043-5323
 DT Journal
 LA German
 CC 1-6 (Pharmacology)
 AB The cytotoxic effect (CE) of various drug combinations was evaluated with the human tumor cell line HT-29. ***Trimidox*** was combined with Ara-C or 2,2'-difluoro-deoxycytidine (DFDC). Synergistic CEs were obsd. Colony area decreased to 71-73% of the control; additive cytotoxicity by sequential treatment with ***Trimidox*** and Ara-C, or by simultaneous treatment with ***Trimidox*** and DFDC. The combination of tiazofurin with allopurinol led to inhibition of hypoxanthine-guanine phosphoribosyl transferase and synergistic CE. Thus, colony area decreased to 69 and 27% of the control; additive cytotoxicity.
- ST cancer chemotherapy colon tumor cell
 IT Drug interactions (synergistic cytotoxic effects of chemotherapeutic drugs on colon tumor cells)
 IT Neoplasm inhibitors (colon, synergistic cytotoxic effects of chemotherapeutic drugs on colon tumor cells)
 IT Intestine, neoplasm (colon, inhibitors, synergistic cytotoxic effects of chemotherapeutic drugs on colon tumor cells)
 IT 9016-12-0, Hypoxanthine-guanine phosphoribosyl transferase
 RL: BSU (Biological study, unclassified); BIOL (Biological study)
 (in synergistic cytotoxic effects of chemotherapeutic drugs on colon tumor cells)
 IT 68-94-0, Hypoxanthine 147-94-4, Ara-C 315-30-0, Allopurinol 60084-10-8, Tiazofurin 95933-74-7, ***Trimidox*** 103882-84-4, 2,2'-Difluorodeoxycytidine
 RL: BAC (Biological activity or effector, except adverse); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
 (synergistic cytotoxic effects of chemotherapeutic drugs on colon tumor cells)
- L3 ANSWER 22 OF 23 HCAPLUS COPYRIGHT 2000 ACS
 AN 1994:569910 HCAPLUS
 DN 121:169910
 TI Biochemical and antitumor activity of ***Trimidox***, a new inhibitor of ribonucleotide reductase
 AU Szekeres, Thomas; Gharehbaghi, Kamran; Fritzer, Monika; Woody, Michael; Srivastava, Arun; van't Riet, Bart; Jayaram, Hiremagalur N.; Elford, Howard L.
 CS Inst. Med. Chem., Univ. Vienna, Austria
 SO Cancer Chemother. Pharmacol. (1994), 34(1), 63-6
 CODEN: CCPHMD; ISSN: 0344-5704
 DT Journal
 LA English
 CC 1-6 (Pharmacology)
 AB ***Trimidox*** (3,4,5-trihydroxybenzamidoxime), a newly synthesized analog of didox (N,3,4-trihydroxybenzamido) reduced the activity of ribonucleotide reductase (EC 1.17.4.1) in exts. of L1210 cells with an IC50 of 5 .mu.M, whereas hydroxyurea, the only ribonucleotide reductase inhibitor in clin. use, exhibited an IC50 of 500 .mu.M. Ribonucleotide reductase activity was also measured in situ by incubating L1210 cells for 24 h with ***Trimidox*** at 7.5 .mu.M (a concn. that inhibited cell proliferation by 50%) or at 100 .mu.M for 2 h; these concns. resulted in a decrease in enzyme activity to 22% and 50%, resp., of the control value. ***Trimidox*** and hydroxyurea were cytotoxic to L1210 cells, with IC50 values of 7.5 and 50 .mu.M, resp. Vs. ribonucleotide reductase, ***Trimidox*** and hydroxyurea had IC50 values of 12 and 87 .mu.M, resp. ***Trimidox*** concn.-dependently increased the life span of mice bearing L1210 leukemia. The antitumor activity appeared more pronounced in female mice than in male mice. These findings suggest that ***Trimidox*** is a new and potent inhibitor of ribonucleotide reductase and that it is a promising candidate for the chemotherapy of cancer in humans.
- ST ***Trimidox*** antitumor ribonucleotide reductase
 IT Neoplasm inhibitors (***Trimidox*** as, ribonucleotide reductase inhibition in relation to)
 IT 127-07-1, Hydroxyurea 69839-83-4, Didox 95933-74-7, ***Trimidox*** (ribonucleotide reductase- and neoplasm-inhibiting activities of)
 IT 9040-57-7 (***Trimidox*** inhibition of, neoplasm inhibition in relation to)
- L3 ANSWER 23 OF 23 HCAPLUS COPYRIGHT 2000 ACS
 AN 1993:182829 HCAPLUS
 DN 118:182829
 TI Prevention of activation of HIV-1 by antiviral agents in OM-10.1 cells
 AU Feorino, P. M.; Butera, S. T.; Folks, T. M.; Schinazi, R. F.
 CS Sch. Med., Emory Univ., Atlanta, GA, 30322, USA
 SO Antiviral Chem. Chemother. (1993), 4(1), 55-63
 CODEN: ACCHEH; ISSN: 0956-3202
 DT Journal
 LA English
 CC 1-5 (Pharmacology)
 AB The development of a reliable and simple system for evaluating compds.

that could prevent activation of latent HIV would allow us to devise new therapeutic approaches. These compds. could eventually be used in combination with drugs that are effective against acute and chronic infections. The OM-10.1 cell line is a chronically infected clone which remains CD4+ until HIV-1 activation with tumor necrosis factor- α . A variety of compds. are known to have antiviral properties against either acutely or chronically infected cells were evaluated for their ability to inhibit HIV induced expression in these cells. The authors also examd. the effect of several compds. that interact with biochem. pathways that may interfere with or enhance the reactivation process. These included nucleoside analogs, cytokines, steroidal and non-steroidal anti-inflammatory agents, polyoxometalates, a TAT inhibitor, various natural products (including nerve growth factor, N-acetyl-L-cysteine, taxol, and interferons), TIBO, porphyrins, and various oligomers. CD4 cellular expression and supernatant reverse transcriptase activity were quantitated as markers of induced viral expression. Among several compds. evaluated, 3'-fluoro-3'-deoxythymidine (FLT), interferon γ , Ro 5-3335 (a TAT inhibitor) and desferrioxamine were modest and selective inhibitors of HIV-1 activation.

ST antiviral HIV1 activation inhibition; immunodeficiency virus activation antiviral

IT Virucides and Virustats

Interferons

(HIV-1 activation prevention by)

IT Virus, animal

(human immunodeficiency 1, activation of, antiviral agents in prevention of)

IT 50-24-8, Prednisolone 50-78-2, Aspirin 53-43-0, Dehydroepiandrosterone 53-86-1, Indomethacin 61-68-7, Mefenamic acid 70-51-9, Desferrioxamine 127-07-1, Hydroxyurea 616-91-1, N-Acetyl-L-cysteine 651-48-9 3056-17-5, 2',3'-Didehydro-3'-deoxythymidine 4428-95-9 7481-89-2, 2',3'-Dideoxycytidine 9061-61-4, Nerve growth factor 15687-27-1, Ibuprofen 25526-93-6, 3'-Fluoro-3'-deoxythymidine 25609-92-1 25609-92-1D, thiolated 28507-02-0 28802-05-3 30195-30-3, Ro 5-3335 30516-87-1, 3'-Azido-3'-deoxythymidine 30811-80-4 33069-62-4, Taxol 33369-31-2, Zomepirac 35218-75-8 35711-34-3, Tolactin 36322-90-4, Piroxicam 38194-50-2, Sulindac 41107-56-6, 3'-Fluoro-2',3'-dideoxyuridine 51246-79-8, 3'-Fluoro-2',3'-dideoxycytidine 69655-05-6, 2',3'-Dideoxyinosine 69839-83-4, Didox 81777-50-6 84472-85-5, CS-87 87190-79-2, CS-92 89899-81-0, HPA-23 92739-63-4 95933-74-7, ***Trimidox*** 115249-95-1, 3'-Fluoro-2',3'-dideoxy-5-methylcytidine 120947-28-6, GLQ 223 123027-56-5 126320-77-2, R-82150 132885-30-4 136632-04-7 136632-06-9 136632-07-0 136891-12-8, BCH 189 142168-25-0 142168-26-1 143491-54-7 143823-92-1, JM 2820 (HIV-1 activation prevention by)

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